Application Note No. 2035. Rev. 1.3
NucleoCounter® NC-202™

Count & Viability of Chicken Embryo Fibroblasts - Via2-Cassette™

Product description
The NucleoCounter® NC-202™ automated cell counter and NC-View™ software perform cell counting and viability analyses on a broad range of eukaryotic cells.

Introduction
Easily load a sample from a cell suspension into the Via2-Cassette™ tip by pressing the piston. Inside, cells are stained by two dyes: acridine orange (AO) and DAPI, which label the total and the non-viable cell populations, respectively. Once loaded, place the Via2-Cassette™ in the NucleoCounter® NC-202™ and press RUN to acquire data. The NC-View™ software automatically analyses and presents cell concentration and viability for fast and easy data acquisition.

Application
This application note describes how to determine the cell concentration and viability of cultivated primary chicken embryo fibroblasts (CEF). The Via2-Cassette™ provides a simple and robust method to determine cell concentration and viability with the NucleoCounter® NC-202™.

Procedure
Harvest ≥ 200 µl of cultivated primary chicken embryo fibroblasts cell sample, preferably in the optimal culture medium for the specific cell type. Transfer a representative sample to a 1.5 ml micro centrifuge tube from which an aliquot can be drawn using the Via2-Cassette™.

Materials needed
• CEF sample in suspension
• Via2-Cassette™

Procedure
1. Mix the cell suspension to homogeneity
2. Load a cell sample by inserting the tip of the Via2-Cassette™ in the cell suspension, then press the piston
3. Insert the loaded Via2-Cassette™ in the NucleoCounter® NC-202™, select the ‘CEF’ protocol and press RUN

After approximately 30 seconds, the cell concentration and viability of the sample are displayed. The results available are: Total (cells/ml), Live (cells/ml), Dead (cells/ml), RBC (cells/ml; Red Blood Cells), Viability (%), Aggregates (%; cell clumps of 5 cells or more), DebrisIndex™, Dilution factor and Status.
Notes
To ensure robust and reliable results, the cell suspension concentration should be in the range of 5×10⁴ - 5×10⁶ cells/ml. If the cell concentration is above 5×10⁶ cells/ml, dilute it with growth medium to ensure accurate cell counting. The diluted cell sample is then counted as described above.

Viability
The viability percentage is calculated as follows:

\[ \% \text{ Viability} = \frac{C_t - C_{nv}}{C_t} \times 100\% \]

% Viability: The percentage of viable cell in the cell sample
C\text{t}: The total concentration of cells (i.e. acridine orange positive cells)
C\text{nv}: The concentration of non-viable cells (i.e. DAPI positive cells)

Picture of NC-View™ software after running a CEF protocol on a CEF cell sample using the NucleoCounter® NC-202™. Acridine orange (AO) and DAPI channels are shown in green and blue, respectively. Enabling the image overlay displays live cells (yellow), dead cells (red), and red blood cells (RBCs; green) as identified by the software. Counting results are presented in the right panel and in the file list below.
Troubleshooting

Inaccurate cell count: My cell count is either too high or low
When analyzing a new cell line, it is important to verify that the cells are correctly identified and recorded. Cells identified by the NC-View™ software can be shown by clicking cell overlay, right panel (see figure). All cells should be highlighted, while cellular debris should be excluded.

Imprecise cell count: I see large variation between technical replicates
The cell counting precision, often quantified as the coefficient of variation from replicate counts, is affected by many variables, including:

2. Liquid handling: The cell suspension should be thoroughly mixed before the sample is aspired into the Via2-Cassette™
3. Cell sample size: The Via2-Cassette™ can load from 200 µl sample in a 1.5 ml tube, however increasing the sample volume improves the precision
4. Consistent protocol execution: Human variation and possibly error in sample handling causes variation between samples and replicas
5. Sample preparation: Ensure that cell sampling and sample dilutions are made using wide orifice tips to avoid 'bottleneck effects'

Handling and storage
For handling and storage of ChemoMetec® instruments and reagents, cassettes and NC-Slides refer to the corresponding product documentation. For other reagents refer to the material data sheet from the manufacturer of the reagents and chemicals.

Warnings and precautions
For safe handling and disposal of the ChemoMetec® reagents, cassettes and NC-slides refer to the corresponding product documentation and the NucleoCounter® NC-202™ user guide. For other reagents refer to the safety data sheet from the manufacturer of the reagents and chemicals required for this protocol. Wear suitable eye protection and protective clothes and gloves when handling biologically active materials.

Limitations
The NucleoCounter® NC-202™ system is FOR RESEARCH USE ONLY. NOT FOR DIAGNOSTIC OR THERAPEUTIC USE. The results presented by the NucleoCounter® NC-202™ system depend on correct use of the reagents, Cassettes and the NucleoCounter® NC-202™ instrument and might depend on the type of cells being analyzed. Refer to the NucleoCounter® NC-202™ user’s guide for instructions and limitations.

Liability disclaimer
This application note is for RESEARCH PURPOSES ONLY. It is not intended for food, drug, household, or cosmetic use. Its use must be supervised by a technically qualified individual experienced in handling potentially hazardous chemicals. The above information is correct to the best of our knowledge. Users should make independent decisions regarding completeness of the information based on all sources available. ChemoMetec A/S shall not be held liable for any damage resulting from handling or contact with the above product.

Product disclaimer
ChemoMetec A/S reserves the right to introduce changes in the product to incorporate new technology. This application note is subject to change without notice.

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